

AMENDMENTS TO THE CLAIMS

This listing of claims will replace all prior versions and listings of claims in the application:

LISTING OF CLAIMS:

1.-19. **(Canceled).**

20. **(Currently Amended)** A method for identifying a candidate protein useful as an anti-infective, comprising:

(a) calculating computationally protein sequence-based attributes from protein sequences of a pathogenic organism, wherein said protein sequences are predicted from whole genomic sequences or are predicted from partial genomic sequences comprising at least one chromosome, and wherein said protein sequence-based attributes comprise~~are selected from a group consisting~~: percentage of charged amino acids, percentage hydrophobicity, distance of protein sequence from a fixed reference frame, measure of dipeptide complexity, and measure of hydrophobicity from a fixed reference frame;

(b) clustering computationally said protein sequences based on said protein sequence-based attributes using Principle Component Analysis, and displaying the results of said clustering;

(c) identifying computationally outlier proteins, wherein said outlier proteins appear outside a main cluster;

(d) comparing outlier proteins with protein sequences in databases;

(e) selecting an outlier protein for further testing as an anti-infective;

and

(f) validating said outlier protein as an anti-infective.

21. **(Previously presented)** The method of claim 20, wherein said pathogenic organism is selected from the group consisting of *B.burgdorfei*, *C.jejuni*, *C.pneumoniae*, *C.trachomatis*, *H.influenzae*, *H.pylori*, *L.major*, *M.genitalium*, *M.pneumoniae*, *M.tuberculosis*, *N.meningitis*, *P.aeruginosa*, *P.falciparum*, *R.prowazekii*, *T.pallidum*, and *V.cholerae*.

22. **(Currently amended)** The method of claim 20, wherein said protein sequence-based attributes ~~comprise~~are selected from the group consisting of fixed protein attributes and variable protein attributes.

23. **(Previously presented)** The method of claim 22, wherein a variable protein attribute is a distance of protein sequence from a variable reference frame.

24. **(Previously presented)** The method of claim 20, wherein said clustering is done by Principle Component Analysis using correlation coefficient between said protein sequence-based attributes.

25. **(Canceled)**

26. **(Previously presented)** The method of claim 20, wherein said outlier protein is non-homologous to known anti-infective proteins from a pathogen selected from the group consisting of

B.burgdorfei, *C.jejuni*, *C.pneumoniae*, *C.trachomatis*, *H.influenzae*, *H.pylori*, *L.major*, *M.genitalium*,
M.pneumoniae, *M.tuberculosis*, *N.meningitis*, *P.aeruginosa*, *P.falciparum*, *R.prowazekii*, *T.pallidum*, and
V.cholerae.

27. **(Previously presented)** The method of claim 20, wherein said outlier protein has an amino acid sequence selected from the group consisting of SEQ ID Nos: 1-31.

28. **(Previously presented)** The method of claim 20, wherein said outlier protein has an amino acid sequence selected from the group consisting of SEQ ID Nos: 32-118.

29. **(Currently Amended)** The method of claim 20, wherein steps are performed by a computer system comprising:

(1) a central processing unit (CPU), wherein said CPU executes a ~~DISTANCE~~ program that calculates protein sequence-based attributes, wherein said protein sequence-based attributes comprise: percentage of charged amino acids, percentage hydrophobicity, distance of protein sequence from a fixed reference frame, measure of dipeptide complexity, and measure of hydrophobicity from a fixed reference frame; and clusters protein sequences based on said protein sequence-based attributes using Principle Component Analysis, thereby producing results;

(2) a memory device accessed by said CPU, wherein said memory device stores said results;

(3) a display on which said CPU displays said results in response to user inputs; and

(4) a user interface device.

30. **(Previously presented)** The method of claim 20, further comprising using said outlier protein for a diagnostic purpose.

31. **(Canceled)**

32. **(Previously presented)** The method of claim 20, further comprising using said outlier protein for a therapeutic purpose.

33. **(Previously presented)** The method as of claim 20, wherein said outlier protein can elicit an immune response.